REMARKS

Claims 1 to 8 are pending in the application.

The purpose of this amendment is to insert the reference to the parent application of which this is a continuation, to place the application headings in appropriate U.S. form, to revise the specification and claims to proper U.S. form and correct syntax and grammar, and to delete multiple dependencies in the claims. Such amendments are formal in nature and no new matter is added by any of the above amendments. Entry of this amendment and early examination of this application are respectfully solicited.

Respectfully submitted,

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November 24, 2003 By: Mil

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WWS:srn Enclosure

(BX2390M)

Maked-up Version of Substitute Specification

<u>METHOD FOR MEASURING THE CONCENTRATION OF WATER IN ARGON,</u>

<u>HYDROGEN, NITROGEN AND HELIUM BY MEANS OF IONIZATION MOBILITY</u>

SPECTROMETRY

TITLE OF THE INVENTION

5 [0001] Method for Measuring the Concentration of Water in Argon, Hydrogen, Nitrogen and Helium by Ionization Mobility Spectrometry

CROSS-REFERENCE TO RELATED APPLICATIONS

[0002] This application is a continuation of International Application No. PCT/IT02/00370, filed June 6, 2002, which was published in the English language on December 12, 2002, under International Publication No. WO 02/099405 A2 and the disclosure of which is incorporated herein by reference.

BACKGROUND OF THE INVENTION

- [0003] The present invention relates to a method for measuring the concentration of water in argon, hydrogen, nitrogen and helium by means of ionization mobility spectrometry.
- 15 [0004] These gases are widely used in the semiconductor industry as transport gases in which reactive species are diluted or as support gases for plasma formation in the cathodic deposition processes (in particular, helium and argon are employed for these uses), as well as real reagents in the process (in particular in the case of nitrogen). Among these gases, argon is the most important for the industry: in. In the rest of this textspecification this gas will be mainly referred to, but the invention may be also be applied with the same results to the other cited gases.
 - [0005] The pureness of the argon employed in the semiconductor industry is particularly important; as. As a matter of fact, contaminants which may be present in the reagents or in the reaction environment can be incorporated into the solid state devices, thereby altering their electrical or magnetic properties and thus leading to production rejects.
- 25 [0006] The argonArgon purification is the subject-matter of various patents, such as for instanceBritish Patent GB-B-2177079 (similarly, British Patent GB-B-2177080 relates to nitrogen purification and US 5558844U.S. Patents 5,558,844 and 55566035,556,603 relate to hydrogen purification). According to this patent, argon is purified by passing it through a bed made of a getter material (an alloy based on zirconium, vanadium and iron) kept at a temperature comprised between 350 and 450 °C; alternatively. Alternatively, purifiers working at room temperature are commonly employed, which are based on the use of nickel generally dispersed onto a high surface area support,

such as for instance alumina or molecular sieves. With these methods the impurity content can be reduced below one part per billion (ppb, equivalent to one impurity molecule for every 10⁹ molecules of argon).

[0007] In Under these conditions it is also necessary to allow the for control of the gas purity and its constancy in over time, for detecting increments of the impurity concentration, due for example to working anomalies of the purifiers, tightness losses of the gas lines or elseother reasons.

[0008] A particularly interesting technique for carrying out this analysis is the-ionization mobility spectrometry, also referred to in the field with the initials IMS (the same initials are also used for the instrument carrying out this technique, i.e. the, "Ionization Mobility Spectrometer").

The interest for Interest in this technique derives from its very high sensibilitysensitivity, combined with the limited size and costs of the instrument; by. By operating inunder suitable conditions species in the gas or vapor phase can be detected in a gas medium in quantities of picograms (pg, i.e., 10^{-12} grams), or in concentrations of parts per trillion (ppt, equivalent to a molecule of the analyzed substance for every 10^{12} molecules of the gas sample). IMS instruments and analysis methods in which these are employed are disclosed, for example, in US patents 5457316 and 5955886U.S. Patents 5,457,316 and 5,955,886 in the name of the US company PCP Inc.

[0009] An IMS instrument is essentially made up of a reaction zone, a separation zone and a charged particles detector.

[0010] In the reaction zone the ionization of the sample comprising the gases or vapors to be analyzed in a transport gas takes place, commonly by means of beta-radiation emitted by ⁶³Ni. The ionization takes place mainly on the transport gas with the formation of the so-called "reagent ions", the," whose charge of which is then distributed on the present-species present according to their electronic or proton affinities or to their ionization potentials. The book "Ion Mobility Spectrometry" by G. A. Eiceman and Z. Karpas, published in 1994 by CRC Press, can be referred to for an illustration of the (rather complex) charge transfer principles which are the basebasis of the ionization mobility spectrometry technique.

[0011] The reaction zone is divided from the separation zone by a grid which, kept at a suitable voltage, prevents the ions produced in the reaction zone from entering into the separation zone. The moment inat which the grid voltage is turned off, thus allowing the ions to enter the separation zone, is the "time zero" of the analysis. The separation zone comprises a series of electrodes, which create an electric field such that the ions are carried from the reaction zone towardstoward the detector. This zone is kept at atmospheric pressure: therefore. Therefore, the motion speed of the ions depends on the electric field and on the cross-section thereof in the gaseous medium. By recording

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the current reading of the particle detector according to the time elapsed from the "time zero"," peaks corresponding to the so-called "drift time" of the different present ions are obtained; by. By determining the drift time it is possible to go back to the presence of the substances which are the object of the analysis.

In spite of its conceptual simplicity, the application of the technique involves some difficulties in the interpretation of the analysis results.

[0013] The instrument, analogously to chromatographs, provides as a result of the analysis the crossing times (drift times in the case of the IMS) of the ions corresponding to the present-species present, but it does not provide indications about the chemical nature of the ion corresponding to each peak.

[0014] For attributing each peak to a specific ion, the IMS may be connected to a mass spectrometer, which determinates determines the chemical nature of each ion, but in this way the above mentioned advantages of low cost and compactness are renounced.

[0015] Alternatively, it is possible to resort to calibration tests, wherein a sample formed of an extremely pure transport gas containing the substance which is the object of the analysis is used, and the drift time of this latter gas is determined. The However, the analysis in under real conditions is however-complicated, since the various ionic species which are present may lead to phenomena of charge transfer with each other or with present-neutral molecules present, so that the determined drift times can be the characteristic times of species different from those, the whose presence of which has is to be determined.

[0016] A possible method for overcoming the problems found in the real analyses consists in adding the sample gas with a specific substance, called a "doping gas"," which, according to various mechanisms, obtains the effect of notablysignificantly decreasing the sensibilitysensitivity of the measure towardsmeasurement toward the species differing from the one which is the object of the analysis. As examples of practical application of the method of the doping gas may be mentioned US 4551624 U.S. Patent 4,551,624, relating to the addition of ketones or halogenated gases to the gas to be analyzed, US 5032721 and US 5095206; U.S. Patents 5,032,721 and 5,095,206, relating, respectively, to the use of phenols and sulfur dioxide in the analysis of acid gases; and US 5238199, U.S. Patent 5,238,199, relating to the use of amines in the analysis of chlorine dioxide.

However, the doping gas method disadvantageously requires that a tank of this gas and means for its dosage in the gaseous medium are added to the system, thus leading to a complication of the measuring system based on the IMS instrument.

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[0017] In the methods not based on the employment of a doping gas the possibility of carrying out a quantitative analysis is bound to the presence of a reactant ion. As previously described, the reactant ion generally is an ion corresponding to the gas present in higher amount in the mixture. Then, the reactant ion formed in the ionization zone transfers the charge to the other present species present according to complex chemical balances. When the impurities concentration of impurities increases, also the charge quantity transferred thereto from the reactant ion also increases, until the latter runs out is extinguished. In the IMS spectrum this mechanism is reflected by the intensity increase of the peaks relevant related to the impurities and by the simultaneous intensity decrease of the peak of the reactant ion, commonly defined in the field as "Reactant Ion Peak" or RIP, up to its extinction. Obviously, when this condition is reached, the concentration of the ions relating to the impurities and the intensity of the relevant peaks in the spectrum do not grow any more-also, even if the effective impurity concentration increases and therefore it is not no longer possible any more to carry out a quantitative IMS analysis in this way. In the presence of water in argon, the RIP runs out is extinguished with concentrations of about 10-15 ppb; thus, at. Thus, according to the state of the art, this value is the maximum measuremeasurement limit of this impurity in argon with the IMS technique.

BRIEF SUMMARY OF THE INVENTION

1. It is therefore an object of the present invention to provide a method which is free from saidthese disadvantages, i.e., a method allowing to measurethe measurement of water concentrations greater than 15 ppb. This object is achieved with a method, the main features of which are disclosed in the first claim and other features are disclosed in the subsequent by a method for measuring the concentration of water in argon, hydrogen, nitrogen and helium by ionization mobility spectrometry (IMS), comprising the followings operative steps:

introducing a gas mixture to be analyzed into an IMS instrument (10) with a counterflow of pure gas;

obtaining a signal (19) variable over time and proportional to a number of ions detected by an ion detector (14) of the IMS instrument (10);

determining two time intervals (A, B) corresponding to drift times in the IMS instrument (10) of H_3O^+ and $(H_2O)_2^+$ ions present in the gas mixture;

obtaining peaks of the signal (19) in the two determined time intervals (A, B); and

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calculating the water concentration in the gas mixture according to a ratio of intensity of the two peaks obtained in the signal (19). Other features of the method are disclosed below and in the dependent claims.

[0018] Thanks to the calculation of the ratio between the intensities of the peaks detected at determined time positions in the signal coming from an IMS instrument, the method according to the present invention allows to precisely measure precise measurement of water concentrations in argon up to at least 30 ppb; the. The measurement of the peak intensity in the practice generally consists intoin the measurement of their area.

[0019] Furthermore, the method according to the present invention, when applied to the ultrapure argon employed in the semiconductor industry (such as hydrogen, nitrogen and helium), does not require either doping gases or gas purification processes before the IMS analysis, except when the same gas has impurities in relatively high concentrations, generally greater than about 10 ppb for each impurity.

[0020] According to a particular aspect of the invention, the calculation of the water concentration can be carried out in an automatic manner by using a logarithmic formula which can be easily implemented in <u>an electronic ealculation meanscalculator</u>, for instance in a personal computer.

Further advantages and features of the method according to the present invention will become clear to those skilled in the art from the following detailed and non-limiting description of an embodiment thereof with reference to the attached drawings, wherein:

BRIEF DESCRIPTION OF THE SEVERAL VIEWS OF THE DRAWINGS

[0021] The foregoing summary, as well as the following detailed description of the invention, will be better understood when read in conjunction with the appended drawings. For the purpose of illustrating the invention, there are shown in the drawings embodiments which are presently preferred. It should be understood, however, that the invention is not limited to the precise arrangements and instrumentalities shown. In the drawings:

[0022] figure Fig. 1 shows is a schematic view diagram of the an instrument for carrying out the method according to the present invention;

[0023] figure Fig. 2 shows is a graph showing some spectrums spectra obtained through the instrument of figure Fig. 1; and

[0024] figure Fig. 3 shows is a graph of some test diagrams curves of the instrument of figure Fig. 1.

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DETAILED DESCRIPTION OF THE INVENTION

Referring to figure Fig. 1, it is seen that an IMS instrument 10 comprises in a known way a reaction zone 11 and a separation zone 12, which are divided by a grid 13. The gases let into the reaction zone 11 are ionized by means of a radiation source (not shown in the figure Fig.), for instance ⁶³Ni. The so produced ions are guided into the separation zone 12 where, by means of electrodes (not shown in the figure Fig.) suitably arranged along the walls of the samethis zone, the ions are accelerated toward an ion detector 14. The ions motion in the separation zone 12 is schematized in the figure schematically represented in Fig. 1 by the arrows. The gas to be analyzed is let into the IMS instrument 10 through an inlet duct 15. The motion speed toward the ion detector 14 is slowed down by the presence of a counter-current flow of pure argon in the zone 12. For this purpose the IMS instrument 10 comprises at least one duct 16 for introducing argon ininto the separation zone 12 from the side of detector 14. The latter detector 14 is connected to a converter 1717, which transforms the progress during the over time of the electric signal of detector 14 into an analog or digital signal, which in turn can be displayed in turn on a display and/or a printer 18 by means of a spectrum 19 representing the result of the analysis.

[0026] Ducts 15 and 16 are preferably made upmanufactured according the usual techniques of the pure gas technology, which generally involve the use of electropolished electro-polished steel pipes for ensuring extremely limited degassing from the internal walls of the samethese ducts.

of the IMS instrument 10 and analyzed according the typical way of this technique, i.e., by employing in the separation zone 12 of the IMS instrument 10 a counter-flow of a gas which does not interfere with the measurement, in particular argon; as it. As will be described in detail later on, the position in the spectrum (in milliseconds, ms) of the peak due to argon, i.e., of the RIP, is known by the by knowledge of the test parameters; when. When the measuring system detects the RIP extinction, two peaks corresponding to two different ionic species relating to water are sought; by. By comparing the intensities of these two peaks, the water concentration in argon is calculated, in particular by means of a logarithmic formula.

<u>[0028]</u> The position of the RIP, seas well as any other peak in an IMS spectrum, depends on the physical parameters with which the analysis is carried out. In particular, the most important parameter is the temperature of the gas in the separation zone: <u>Thus</u>, considerable temperature changes with respect to a reference temperature may change the ratios between the time position of various peaks in the spectrum, thereby making the analysis impossible. A second important parameter is the ratio between the applied electric field and the gas pressure in the separation zone;

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ehanges. Changes of this ratio cause proportional changes of the positions of all the peaks in the spectrum, so that the spectrum is "compressed" or "expanded"," although the ratios between of the various peak positions are keptmaintained. Anyway, for convenience, it is preferable that the drift times remain fixed, for facilitating the identification of the different peaks. Finally, the spectrum is influenced by the ratio between the mass flow of gas to be analyzed and the mass flow of the counter-current gas, even if in lower measure with respect tothough to a lesser extent than the previous parameters. The two peaks relating to water, which are important for the method according to the invention, are the peaks corresponding to H₃O⁺ and (H₂O)₂⁺ ions. Therefore, for knowingto know the time position of these two peaks, it is necessary to carry out an IMS guidingcalibration test inunder the same conditions of the real analysis, with a water content in argon such that the RIP is not extinct. A complete spectrum is thus obtained: the The presence of the RIP allows to assignassignment of the other peaks to the corresponding species, this correspondence being keptmaintained in the real analysis (without RIP).

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EXAMPLE

[0029] The invention will be further explained by the following example, relating to a series of tests of water analysis with different concentrations in argon. In all the tests, the gas mixture flowing into the IMS instrument 10 is keptmaintained equal to 0.50.5 liters per minute, and the gas temperature is 110 °C. The sample ionization is carried out by a ⁶³Ni radioactive source. The ions so generated-ions are neutralized onto grid 13 until the voltage of the latter is canceled, thereby allowing their entrance into the separation zone 12. The cancellation time of the grid voltage is 200 microseconds (µs) for each test. The tests are carried out with an IMS instrument 10 wherein the separation zone 12 is 8 cm long; in. In every test the acceleration electric field is equal to 128 V/cm. From preliminary guidingcalibration tests it resulted is determined that inunder these conditions the typical drift times of the species present in the tests are generally eomprised between 15 and 30 milliseconds (ms); moreover, Moreover, with a suitable test it has been determined that in under these conditions the peaks of H₁O⁺ and (H₂O)₂⁺ are comprised between 15, 515.5 and 17 ms and between 17 and 19 ms, respectively (with maximums at about 16 and 47,817.8 ms, respectively). The results of all the tests are reported in spectrums 19 spectra 19 of the display and/or printer 18 (Fig. 1), wherein the peaks have an area proportional to the concentration of a given ion according to the drift time of the same. The peak intensity is expressed in volts (V), while the current directly measured by detector 14 (number of ions colliding onto the detector per time-unit time) is

transformed in volts by converter 17. The counter-flow of pure argon coming from duct 16 has a rate of 2 liters per minute. Generally, the ratio between the flow rate of the gas to be analyzed and the counter-flow of pure argon can vary between 1:10 and 1:1.

<u>[0030]</u> Figure Fig. 2 shows a plurality of overlapping spectrums spectra obtained by analyzing with the IMS instrument 10 a corresponding plurality of mixtures of water, argon and very little impurities, wherein only the water concentration changes. Along the <u>abscissa (x-axis-of-abscissas)</u> is shown the drift time of the ions in milliseconds, while along the <u>ordinate (y-axis-of-ordinates)</u> is shown a value proportional to the intensity of the signal emitted by the ion detector 14, for instance a voltage expressed in volts. The numbers inside the diagram distinguish instead the different spectrums spectra according to the water concentration expressed in ppb present in the gas let into the IMS instrument 10.

[0031] As it can be noticed seen, without the presence of water, i.e., with 0 ppb of water, the relevant spectrum (shown with a bold line) has 4 peaks A, C, D and E, the latter last of which is the tallest. The latter last peak E, which in under the conditions of the present test is in the time interval emprised between 23 and 25 ms, corresponds to the peak of the Ar⁺ ions, i.e., to the RIP, while the other peaks A, C and D, emprised in the time intervals between 15,5 ms and 17 ms, 19 and 21 ms, and 21 and 23 ms, respectively, correspond to the ions of the impurities present in the gas mixture. In particular, the peak in the first time interval A is caused by the ions of H_3O^+ , the second peak C is caused by the ions of the impurities, and the third peak D by the ions of H_2O^+ . The ions of H_2O^+ and H_3O^+ of peaks A and D are due to very small water concentrations which cannot be eliminated from the incoming gas mixture.

[0032] By increasing to 5 ppb the water concentration in the gas mixture and by keeping unchanged the other variables of the test, it can be noticed seen how the heights of peaks C and E notably markedly fall, so as also and the height of peak D is also slightly reduced, while the height of peak A grows. Further, in the time interval between 17 and 19 ms a new peak B which is caused by the $(H_2O)_2^+$ ions can be distinguished. At 5 ppb water, peaks A and B have heights similar to each other, and in any case higher than the remaining peaks C, D, and E.

[0033] By increasing the water concentration in the mixture from 5 ppb to about 30 ppb, the height of peak B grows, while the height of peak A falls and peaks C, D and E substantially disappear. In particular, at around 18 ppb of water the height of peak A is equal to the height of the same peak with 0 ppb of water.

[0034] By further increasing the water concentration in the mixture beyond 30 ppb, the system is saturated and only peak B remains with a constant height.

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[0035] As already said, the positions of peaks A and B were already previously determined with a guidingcalibrating test, thereby resulting in the time intervals between 15,515.5 and 17 ms and between 17 and 19 ms. By evaluating the intensity of these peaks, and in particular their area, the corresponding levels of the signal measured by the ion detector 14 are obtained. The water concentration contained in the analyzed gas can be determined by comparing saidthe levels. In particular, for water concentrations lower than 30 ppb, it has been discovered that saidthe comparison can be expressed with the following formula:

> $ppb_{H2O} = K \ln ((HB + HA) / HA)$, wherein: ppb_{H2O} is the water concentration, K is a positive constant, HA is the area of peak A, and HB is the area of peak B.

[0036] FigureFig. 3 shows a graph comprising showing the change during theover time (abscissas abscissa) of two overlapping diagrams curves of the water concentration 15 (ordinates ordinate). The first diagram curve, made up of a continuous stepped line, shows the water concentration effectively present in a reference mixture let into the IMS instrument 10, while the second diagram curve, made up of a dotted line, shows the water concentration concentrations measured by means of the above described method. As it can be noticed seen, the trendcourse of the second diagram curve substantially follows the trend course of the first diagram curve up to 30 ppb. 20 while differs differing beyond this concentration, in at which the instrument is saturated.

[0037] It will be appreciated by those skilled in the art that changes could be made to the embodiments described above without departing from the broad inventive concept thereof. It is understood, therefore, that this invention is not limited to the particular embodiments disclosed, but it is intended to cover modifications within the spirit and scope of the present invention as defined by the appended claims.

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CLAIMS

We claim:

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1. A method for measuring by means of ionization mobility spectrometry thea concentration of water in argon, hydrogen, nitrogen and helium, characterized by ionization mobility spectrometry (IMS), comprising the followings operative steps:

introducing thea gas mixture to be analyzed into an IMS instrument (10) with a counterflow of pure gas;

obtaining a signal (19) variable <u>during theover</u> time and proportional to <u>thea</u> number of ions detected by an ion detector (14) of the IMS instrument (10);

determining two time intervals (A, B) corresponding to the drift times in the IMS instrument (10) of the H_3O^+ and $(H_2O)_2^+$ ions present in the gas mixture;

obtaining the peaks of saidthe signal (19) in the two determined time intervals (A, B); and

calculating the water concentration in the gas mixture according to thea ratio between theof intensity of the two peaks obtained in the signal (19).

2. A<u>The</u> method according to claim 1, eharacterized in that<u>wherein</u> the water concentration in the analyzed gas is calculated by-means of the following formula:

 $ppb_{H2O} = K \ln ((HB + HA) / HA)$, wherein:

ppb_{H2O} is the water concentration in ppb,

K is a positive constant,

HA is the intensity of the peak of the signal (19) in the time interval (A) corresponding to the drift times of the H_3O^+ ions in the IMS instrument (10); and

HB is the intensity of the peak of the signal (19) in the time interval (B) corresponding to the drift times of the $(H_2O)_2^+$ ions in the IMS instrument (10).

- 3. A<u>The</u> method according to claim 2, wherein the area areas of the peaks is are employed as a measure measures of saidthe intensity of the same respective peaks.
- 4. A<u>The</u> method according to claim 2, wherein the heightheights of the peaks is are employed as a measure measures of saidthe intensity of the same respective peaks.
- 5. A<u>The</u> method according to one of the previous claims, characterized in that claim 1, wherein the two time intervals (A, B) corresponding to the drift times of the H₃O⁺ and (H₂O)₂⁺ ions in the IMS instrument (10) are determined with a preliminary guiding test carried out with values of the operative parameters equal to those employed in the an actual analysis.
- 6. A<u>The</u> method according to claim 5, wherein <u>saidthe</u> operative parameters comprise at least thea temperature of the analyzed gas mixture.
 - 7. A<u>The</u> method according to claim 6, wherein saidthe operative parameters <u>further</u> comprise also thean electric field in thea separation zone (12) of the IMS instrument (10).
- 8. AThe method according to ene of claims 5 to 7, characterized in that claim 5, wherein the analysis is carried out with gases at the a temperature of 110 °C and with an electric field of 128

 V/cm in the a separation zone (12) of 128 V/emthe IMS instrument (10), and the two time intervals (A, B) corresponding to the drift times of the H₃O⁺ and (H₂O)₂⁺ ions in the IMS instrument (10) are comprised between 15,515.5 and 17 ms (A) and between 17 and 19 ms (B).

ABSTRACT OF THE DISCLOSURE

A method for is provided for measuring by means of ionization mobility spectrometry relatively high concentrations of water in argon, hydrogen, nitrogen and helium, eharacterized by comprising including the followings operative steps: (a) introducing the gas to be analyzed into an IMS instrument (10) with a counter-flow of pure gas; (b) obtaining a signal (19) variable over time and proportional to the number of ions detected by an ion detector (14) of the IMS instrument (10); (c) determining two time intervals (A, B) corresponding to the drift times in the IMS instrument (10) of the H₃O⁺ and (H₂O)₂⁺ ions; (d) obtaining the peaks of the signal (19) in the two determined time intervals (A, B); and (e) calculating the water concentration in the gas to be analyzed according to the ratio between the intensity of the two peaks obtained in the signal (19).

introducing the gas to be analyzed into an IMS instrument (10) with a counter flow of pure gas; obtaining a signal (19) variable during the time and proportional to the number of ions detected by an ion detector (14) of the IMS instrument (10);

determining two time intervals (A. B) corresponding to the drift times in the IMS instrument (10) of the H₂O⁺ and (H₂O)₂⁺ions;

obtaining the peaks of said signal (19) in the two determined time intervals (A, B);

calculating the water concentration in the gas to be analyzed according to the ratio between the intensity of the two peaks obtained in the signal (19).

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